	Туре	L #	Hits	Search Text	DBs	Time Stamp	Comment
1	BRS	L1	69	urinary adj trypsin adj inhibitor	USPA T; US-P GPUB; EPO; DERW ENT	2002/04/0 2 12:43	
2	BRS	L2	24	urine adj trypsin adj inhibitor	USPA T; US-P GPUB; EPO; DERW ENT	2002/04/0 2 12:44	
3	BRS	L4	32772 1	calcium	USPA T; US-P GPUB; EPO; DERW ENT	2002/04/0 2 12:44	
4	BRS	L5 <sub>.</sub>	27	3 and 4 .	USPA T; US-P GPUB; EPO; DERW ENT	2002/04/0 2 12:44	
5	BRS	L6	33	polycarboxylic adj chelat\$	USPA T; US-P GPUB; EPO; DERW ENT	2002/04/0 2 12:45	
6	BRS	L7	1	3 and 6	USPA T; US-P GPUB; EPO; DERW ENT	2002/04/0 2 12:46	

	Туре	L #	Hits	Search Text	DBs	Time Stamp	Comment s
7	BRS	L8	4317	trypsin adj inhibitor\$	USPA T; US-P GPUB; EPO; DERW ENT	2002/04/0 2 12:50	
8	BRS	L9	2	6 and 8	USPA T; US-P GPUB ; EPO; DERW ENT	2002/04/0 2 12:47	
9	BRS	L10	54857	edta or egta	USPA T; US-P GPUB; EPO; DERW ENT	2002/04/0 2 12:48	
10	BRS	L11	2286	8 and 10	USPA T; US-P GPUB; EPO; DERW ENT	2002/04/0 2 12:48	
11	BRS	L12	649	8 same 10	USPA T; US-P GPUB; EPO; DERW ENT	2002/04/0 2 13:04	
12	BRS	L13	2418	8 and substrate\$	USPA T; US-P GPUB ; EPO; DERW ENT	2002/04/0 2 12:49	

	Туре	L #	Hits	Search Text	DBs	Time Stamp	Comment s
13	BRS	L14	684	8 and urine	USPA T; US-P GPUB; EPO; DERW ENT	2002/04/0 2 12:51	
14	BRS	L15	613	14 and buffer	USPA T; US-P GPUB ; EPO; DERW ENT	2002/04/0 2 12:51	
15	BRS	L3	69	urinary adj trypsin adj inhibitors	USPA T; US-P GPUB ; EPO; DERW ENT	2002/04/0 2 13:03	
16	BRS	L16	2	6 same 8	USPA T; US-P GPUB; EPO; DERW ENT	2002/04/0 2 13:03	
17	BRS	L17	415	4 and 12	USPA T; US-P GPUB; EPO; DERW ENT	2002/04/0 2 13:04	
18	BRS	L18	39	4 same 12	USPA T; US-P GPUB; EPO; DERW ENT	2002/04/0 2 13:04	

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15994 S PROTEASE (W) HIBITOR#
        561554 S CALCIUM
L3
          630 S L1 AND L2
        223837 S URIN?
L4
           16 S L3 AND L4
L6
          329 S URINARY (W) TRYPSIN (W) INHIBITOR#
           16 S L6 AND SUBSTRATE#
L7
L8
        83181 S EDTA OR EGTA
          547 S L1 AND L8
L9
            8 S L9 AND URINE
L10
     2394793 S DETECT OR DETERMINE OR MEASUR?
L11
L12
          53 S L6 AND L11
           21 S L6 (S) L11
L13
         1430 S L1 AND L11
L14
L15
          219 S L1 (S) L11
        83181 S EDTA OR EGTA
L16
       561554 S CALCIUM
L17
L18
         3135 S L16 (S) L17
       80709 S EFFECT (S) CALCIUM
L19
        3425 S L16 AND L19
L20
L21
          159 S OFFSET (S) CALCIUM
L22
           2 S L21 AND L16
L23
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L24
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       107012 CHELAT?
L28
           44 POLYCARBOXYL? (W) CHELAT?
=> s 125 and 128
            0 L25 AND L28
=> s 124 and 128
L30
            1 L24 AND L28
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	15994 S F	2R0	OTEASE (W) I BITOR#
L2			CALCIUM
L3	630	S	L1 AND L2
L4	223837	S	URIN?
L5	16	S	L3 AND L4
L6	329	S	URINARY (W) TRYPSIN (W) INHIBITOR#
L7	16	S	L6 AND SUBSTRATE#
L8	83181	S	EDTA OR EGTA
L9	547	S	L1 AND L8
L10	8	S	L9 AND URINE
L11	2394793	S	DETECT OR DETERMINE OR MEASUR?
L12	53	S	L6 AND L11
L13	21	S	L6 (S) L11
L14	1430	S	L1 AND L11
L15	219	S	L1 (S) L11

A method and a kit for assay urinary trypsin inhibitor

> An accurate method is described for assaying urinary trypsin inhibitor (UTI) by inactivating .alpha.1-antitrypsin (.alpha.1-AT) in a sample, mixing a trypsin soln. with the sample, adding a substrate to initiate an enzyme reaction, and then, measuring a change in absorbance. .alpha.1-AT can be inactivated either by adding a protease other than trypsin to the sample soln. and reacting the protease with .alpha.1-AT to form the complex, or by adding an oxidizing agent to the sample. As a protease to inactivate .alpha.1-AT, elastase or subtilisin can be used. As an oxidizing agent to inactivate .alpha.1-AT, sodium iodate, iodine, copper sulfate or iron trichloride can be used. The amt. of UTI in a urine sample was accurately detd. by this method using subtilisin as an example. PCT Int. Appl., 39 pp.

·SO

CODEN: PIXXD2

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L3	630	S	L1 AND L2
L4	223837	S	URIN?
L5	16	S	L3 AND L4
L6	329	S	URINARY (W) TRYPSIN (W) INHIBITOR#
L7	16	S	L6 AND SUBSTRATE#
$^{L8}$	83181	S	EDTA OR EGTA
L9	547	S	L1 AND L8
L10	8	S	L9 AND URINE

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ANSWER 1 OF 16 CAPLUS COPYRI 2002 ACS
AN 2001:864746 CAPLUS
   135:371994
DN
    Preparation of arginine derivatives for assay of trypsin urinary inhibitor
TI
    Corey, Paul F.; Felman, Steven W.; Rehm, Gary E.; Pugia, Michael J.
IN
    Bayer Corporation, USA
PΑ
    Eur. Pat. Appl., 20 pp.
SO
    CODEN: EPXXDW
DT
    Patent
LΑ
    English
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    EP 1157984 A2 20011128 EP 2001-110138 20010504
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                                      US 2001-844816 20010430
NO 2001002307 A 20011116
JP 2002069055 A2 20020308
PRAI US 2000-203999P P 20000515
                                      NO 2001-2307 20010510
                                      JP 2001-139608 20010510
    MARPAT 135:371994
    ANSWER 2 OF 16 CAPLUS COPYRIGHT 2002 ACS
L7
    2001:850805 CAPLUS
AN
    135:368535
DN
ΤI
    Urinary trypsin inhibitor assay containing a
    polycarboxylic chelating agent
    Rehm, Gary B.; Pugia, Michael J.; Corey, Paul F.
IN
    Bayer Corporation, USA
PA
SO
    Eur. Pat. Appl., 9 pp.
    CODEN: EPXXDW
\mathsf{DT}
    Patent
LΑ
    English
FAN.CNT 1
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                                       US 2001-844815 20010430
    US 2001055816 A1 20011227
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                    A 20011116
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    JP 2002014096
                    A2 20020118
                                       JP 2001-142654 20010514
PRAI US 2000-204032P P 20000515
L7
    ANSWER 3 OF 16 CAPLUS COPYRIGHT 2002 ACS
AN
    1999:626367 CAPLUS
DN
    131:239729
TI
    A method and a kit for assaying urinary trypsin
    inhibitor
IN
    Okamoto, Kazuhiro; Fukunaga, Satoshi
PΑ
    Kyoto Daiichi Kagaku Co., Ltd., Japan
SO
    PCT Int. Appl., 39 pp.
    CODEN: PIXXD2
DT
    Patent
LΑ
    Japanese
FAN.CNT 1
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    PATENT NO. KIND DATE
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                                       ______
    WO 9949076
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                   A1 19990930
                                      WO 1999-JP972 19990226
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        RW: AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL,
            PT, SE
    JP 11318493
                    A2 19991124
                                      JP 1999-36909 19990216
    JP 3059433
                    B2 20000704
                  A 19980320
A 19980320
PRAI JP 1998-72712
    JP 1998-72713
RE.CNT 2 THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD
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ANSWER 4 OF 16 CAPLUS COPYRIGHT 2002 ACS
L7
     1999:196406 CAPLUS
AN
     130:219860
DN
    Method and kit for measuring protease inhibitor, particularly
TΤ
     urinary trypsin inhibitor
    Nanbu, Atsuko; Fukunaga, Satoshi
IN
PΑ
     Kyoto Daiichi Kagaku Co., Ltd., Japan
SO
     Eur. Pat. Appl., 14 pp.
     CODEN: EPXXDW
DT
    Patent
LΑ
    English
FAN.CNT 1
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                                       APPLICATION NO. DATE
    PATENT NO.
     ______
                                        _____
    EP 902091 A2 19990317
EP 902091 A3 20010110
                                       EP 1998-306748 19980824
PΙ
    EP 902091
                    A3 20010110
        R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
            IE, SI, LT, LV, FI, RO
    JP 11075896 A2 19990323
US 6130055 A 20001010
                                        JP 1997-234850 19970829
                    A 20001010
                                        US 1998-135915 19980818
PRAI JP 1997-234850 A
                         19970829
    ANSWER 5 OF 16 CAPLUS COPYRIGHT 2002 ACS
T.7
    1998:21476 CAPLUS
AN
    128:99284
DN
TΙ
    A method for measuring the concentration of protease inhibitors, kit for
    use in such a method and method for dissolving a substrate
    Uenoyama, Harumi; Ohshiro, Kyouichi; Nanbu, Atsuko; Fukunaga, Satoshi
IN
PA
    Kabushiki Kaisha Kyoto Daiichi Kagaku, Japan
    Eur. Pat. Appl., 11 pp.
SO
    CODEN: EPXXDW
DT
    Patent
LΑ
    English
FAN.CNT 1
                   KIND DATE
    PATENT NO.
                                       APPLICATION NO. DATE
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    EP 814167 A1 19971229 EP 1997-304313 19970619
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        R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
            IE, SI, LT, LV, FI, RO
    JP 10070997 A2 19980317
                                       JP 1997-156398 19970613
    US 5856117
                    Α
                         19990105
                                       US 1997-879962 19970620
PRAI JP 1996-162163
                         19960621
    JP 1996-166311
                         19960626
L7
    ANSWER 6 OF 16 CAPLUS COPYRIGHT 2002 ACS
AN
    1996:338759 CAPLUS
    125:56283
DN
ТT
    Secretion of a variant of human single-chain urokinase-type plasminogen
    activator without an N-glycosylation site in the methylotrophic yeast,
    Pichia pastoris and characterication of the secreted product
AU
    Tsujikawa, Muneo; Okabayashi, Ken; Morita, Masanori; Tanabe, Toshizumi
CS
    Green Cross Corp., Hirakata, 573, Japan
SO
    Yeast (1996), 12(6), 541-553
    CODEN: YESTE3; ISSN: 0749-503X
DT
    Journal
LA
    English
L7
    ANSWER 7 OF 16 CAPLUS COPYRIGHT 2002 ACS
AN
    1989:571503 CAPLUS
DN
    111:171503
    Anti-fibrinolytic activity and distribution of urinary
TI
    trypsin inhibitor-related substances
ΑU
    Sumi, Hiroyuki; Yoshida, Etsuo; Nakajima, Nobuyoshi; Hamada, Hiroki;
    Mihara, Hisashi
CS
    2nd Dep. Physiol., Miyazaki Med. Coll., Miyazaki, 889-16, Japan
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Ketsueki to Myakkan (1989 20(2), 171-5
SO
     CODEN: KTMYA3; ISSN: 0386-9717
DT
     Journal
     Japanese
LΑ
L7
     ANSWER 8 OF 16 CAPLUS COPYRIGHT 2002 ACS
AN
     1987:628724 CAPLUS
DN
     107:228724
     Effect of urinastatin on coagulation, fibrinolysis, and platelet
TI
     aggregation in vitro and in vivo
     Sakuragawa, Nobuo; Shimotori, Tomoya; Takahashi, Kaoru; Niwa, Masahiro
ΑU
     Cent. Clin. Lab., Toyama Med. Pharm. Univ., Toyama, Japan
CS
     Saishin Igaku (1987), 42(4), 820-30
SO
     CODEN: SAIGAK; ISSN: 0370-8241
DT
     Journal
LΑ
     Japanese
1.7
     ANSWER 9 OF 16 CAPLUS COPYRIGHT 2002 ACS
AN
     1986:104810 CAPLUS
DN
     104:104810
     Studies on fibrinolytic enzyme in human bronchoalveolar lavage fluid
ΤI
ΑU
     Takagi, Ohmi
CS
     Sch. Med., Kinki Univ., Osaka, Japan
     Kinki Daigaku Igaku Zasshi (1985), 10(3), 221-37
SO
     CODEN: KDIZDD; ISSN: 0385-8367
DT
     Journal
     Japanese
LА
L7
     ANSWER 10 OF 16 CAPLUS COPYRIGHT 2002 ACS
     1985:162840 CAPLUS
AN
     102:162840
DN
TI
     Chemical modification of basic amino acid residues in the urinary
     trypsin inhibitor and its effect on trypsin and
     chymotrypsin inhibitory activities
     Tanaka, Yoshiaki; Soeda, Mitsuo; Moriyama, Shigeru; Sasaki, Koji; Maehara,
ΑU
     Susumu; Kawashita, Eizo
CS
     Zeria Pharm. Co. Ltd., Tokyo, 103, Japan
     Igaku to Seibutsugaku (1984), 109(2), 75-8
SO
     CODEN: IGSBAL; ISSN: 0019-1604
DT
     Journal
     Japanese
LA
L7
     ANSWER 11 OF 16 CAPLUS COPYRIGHT 2002 ACS
AN
     1982:158115 CAPLUS
DN
     96:158115
TI
     Low molecular weight trypsin-plasmin inhibitors isolated from papain
     treated urinary trypsin inhibitor
     Sumi, H.; Toki, N.; Takasugi, S.; Maehara, S.; Maruyama, M.; Akazawa, K.;
AII
     Matsuo, O.; Mihara, H.
     Dep. Physiol., Miyazaki Med. Coll., Miyazaki, Japan
CS
     Thromb. Haemostasis (1982), 47(1), 14-18
SO
     CODEN: THHADQ; ISSN: 0340-6245
DT
     Journal
LA
     English
L7
     ANSWER 12 OF 16 CAPLUS COPYRIGHT 2002 ACS
     1979:415786 CAPLUS
AN
DN
     91:15786
TI
     Inhibition of bovine trypsin with human plasma inhibitors
ΑU
     Takada, Akikazu; Fukuda, Shigeji; Takada, Yumiko
     Sch. Med., Hamamatsu Univ., Hamamatsu, 431-31, Japan
CS
     Thromb. Res. (1979), 14(2-3), 413-22
SO
     CODEN: THBRAA; ISSN: 0049-3848
DT
     Journal
LA
     English
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L7

ANSWER 13 OF 16 CAPLUS COPYRIGHT 2002 ACS

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AN
     1978:592891 CAPLUS
DN
     89:192891
     Comparative study of the amidolytic and caseinolytic methods for the
TΙ
     determination of urinary trypsin inhibitor
     Sundaresh, C. S.; Aroor, A. R.; Pattabiraman, T. N.
ΑU
     Dep. Biochem., Kasturba Med. Coll., Manipal, India
CS
     Indian J. Med. Res. (1978), 68(2), 341-7
SO
     CODEN: IJMRAQ; ISSN: 0019-5340
DΤ
     Journal
LΑ
     English
L7
     ANSWER 14 OF 16 CAPLUS COPYRIGHT 2002 ACS
     1978:85138 CAPLUS
AN
DN
     88:85138
ΤI
     Antitryptic property of cancer-related glycoprotein EDC1
ΑU
     Chawla, Rajender K.; Wadsworth, Allan D.; Rudman, Daniel
CS
     Dep. Med., Emory Univ. Sch. Med., Atlanta, Ga., USA
SO
     Cancer Res. (1978), 38(2), 452-7
     CODEN: CNREA8; ISSN: 0008-5472
DT
     Journal
LΑ
     English
L7
     ANSWER 15 OF 16 CAPLUS COPYRIGHT 2002 ACS
     1978:85021 CAPLUS
AN
     88:85021
DN
TI
     Studies on human urinary enzymes and inhibitors. Concentration method and
     characterization
     Sumi, Hiroyuki; Toki, Naotika; Takada, Yumiko; Takada, Akikazu
ΑU
     2nd Dep. Physiol., Hamamatsu Univ. Sch. Med., Hamamatsu, Japan
CS
     J. Biochem. (Tokyo) (1978), 83(1), 141-7
SO
     CODEN: JOBIAO; ISSN: 0021-924X
\mathsf{DT}
     Journal
     English
LΑ
     ANSWER 16 OF 16 CAPLUS COPYRIGHT 2002 ACS
L7
AN
     1965:45206 CAPLUS
DN
     62:45206
OREF 62:8055e-g
     Ester hydrolysis by urokinase
ΤI
     Lorand, L.; Condit, E. V.
ΑU
CS
     Northwestern Univ., Evanston, IL
SO
     Biochemistry (1965), 4(2), 265-70
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     Journal
     English
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Γ8
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=> s 11 and 18
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## => d 110 1-8 ti abs so

- L10 ANSWER 1 OF 8 CAPLUS COPYRIGHT 2002 ACS
- Improved automated LPA assay and methods of detecting cancer TI
- The present invention relates to an improved enzymic diagnostic assay to AΒ detect carcinoma by measuring various lysophospholipids, including lysophosphatidic acid (LPA), in a patient. In a preferred embodiment, this assay measures the human plasma level of LPA in an automated format with a minimal no. of reagents and with reduced incubation periods. The present invention also comprises several addnl. tech. improvements to the current LPA assays disclosed in the prior art.
- SO PCT Int. Appl., 49 pp.

CODEN: PIXXD2

- L10 ANSWER 2 OF 8 CAPLUS COPYRIGHT 2002 ACS
- Evaluation of intestinal permeability and gluten sensitivity in Soft-Coated Wheaten Terriers with familial protein-losing enteropathy, protein-losing nephropathy, or both
- Objective-To evaluate intestinal permeability and gluten sensitivity in a AB family of Soft-Coated Wheaten Terriers (SCWT) affected with protein-losing enteropathy (PLE), protein-losing nephropathy (PLN), or both. Animals-6 affected adult dogs. Procedure-Intestinal biopsy specimens, urine protein-to-creatinine ratio, serum concns. of albumin and globulin, and concn. of .alpha.1-protease inhibitor in feces were evaluated before, during, and  $13~\mathrm{wk}$  after daily administration of  $10~\mathrm{g}$  of gluten for 7 wk. Eosinophils and lymphocytes-plasmacytes were enumerated in intestinal biopsy specimens. Intestinal permeability was evaluated before and during the sixth week of gluten administration via cellobiose-mannitol and chromium-EDTA absorption tests. Results-Serum globulin concn. decreased significantly after prolonged administration of gluten. Although not significant, there was an increase in lymphocytes-plasmacytes and a decrease in eosinophils in intestinal biopsy specimens. Furthermore, these counts were greater than those reported for clin. normal dogs. Gluten administration did not increase intestinal permeability. Conclusions and Clin. Relevance-Daily administration of gluten was assocd. with a significant decrease in serum globulin concn. in SCWT affected with PLE or PLN, but other variables remained unchanged. Although enhanced wheat-gluten sensitivity may be one factor involved in the pathogenesis of PLE or PLN in SCWT, this syndrome does not appear to be the result of a specific sensitivity to gluten. American Journal of Veterinary Research (2000), 61(5), 518-524 SO CODEN: AJVRAH; ISSN: 0002-9645
- L10 ANSWER 3 OF 8 CAPLUS COPYRIGHT 2002 ACS
- TТ Disposable absorbent article having a skin care composition containing an enzyme inhibitor
- AB An absorbent article, at least a portion of which comprises a skin care compn. that comprises an enzyme inhibitor and is at least partially transferred from the article to the skin of a wearer of the article as a result of normal contact, wearer motion and/or body heat is provided. The enzyme inhibitor is transferred to the skin with the skin care compn. and is available at the skin/urine and skin/feces interfaces to inhibit enzymic activity on the skin and to reduce or prevent the occurrence of inflammation. Repeated application of similar treated articles to the wearer's skin provides an available source with which the enzyme inhibitor transfers onto the skin continuously over time and accumulates to provide a proactive defense against harmful enzymes for the treatment and/or prevention of diaper dermatitis. An absorbent article having a topsheet comprising a skin care compn. and an enzyme inhibitor was prepd. The compn. contained acetohydroxamic acid 1, SEFA cottonate 85, and SEFA behenate 15 parts.

SO PCT Int. Appl., 73 pp.

CODEN: PIXXD2

- L10 ANSWER 4 OF 8 CAPLUS COPIRIGHT 2002 ACS
- TI Simultaneous collection of DNA and non-nucleic acid analytes from oral fluids
- AB This invention provides for a rapid and convenient method of simultaneous collection of both genomic and diagnostic information from a single sample on a bibulous pad by differential extn. of the diagnostic information from the genomic information. Samples may be collected from the mouth, rectum, vagina or nose. It is a surprising discovery of this invention that a PCR assay on the contents of the bibulous pad provides results comparable in reliability, specificity, and sensitivity to the best available serum (blood) based assays. The assays of this invention can be used to confirm each other, either by detecting the genomic information leading to the diagnostic information, or by detecting in the genomic information, a predisposition to a disease and confirming the presence of the disease through diagnostic testing.
- SO PCT Int. Appl., 71 pp. CODEN: PIXXD2
- L10 ANSWER 5 OF 8 CAPLUS COPYRIGHT 2002 ACS
- TI Stabilizing formulation for preserving the integrity of proteins present in a body fluid
- AB The present invention provides stabilizing formulations for maintaining and preserving the integrity of proteins and polypeptides present in the body fluid sample obtained ex-vivo and to be evaluated as a test specimen for either clin., therapeutic, or research purposes. The stabilizing formulations may be prepd. alternatively either as a dry, anhyd. mixt. of powders or as an aq. based liq. contg. the dissolved ingredients in admixt. The invention also provides minimalist stabilizing formulations as well as fortified stabilizing formulations which meet specific uses and applications and may be advantageously employed over a wide variety of different time, temp., and severity of conditions.

An enzyme which cleaves the Glu(143)-Leu(144) bond of pro-urokinase was

SO PCT Int. Appl., 56 pp. CODEN: PIXXD2

AΒ

SO

- L10 ANSWER 6 OF 8 CAPLUS COPYRIGHT 2002 ACS
- TI Characterization of a metalloprotease which cleaves with high site-specificity the Glu(143)-Leu(144) bond of urokinase
- partially purified from the conditioned media of cultured human kidney (HEK) cells. The products of the reaction catalyzed by this enzyme were forms of pro-urokinase and urokinase with Leu(144) as the N-terminal residue. The protease was purified using 3 chromatog. steps: (a) S-sepharose; (b) Zn-Sepharose; and (c) gel filtration. The enzyme was a metalloprotease, requiring Ca2+ or Zn2+, and was inhibited by EDTA The activity was not affected by serine or cysteine protease inhibitors. Although the enzyme was purified >1000-fold from the culture media, a prep. of homogeneous protein could not be obtained. Completion of the isolation of the protease will allow detn. of whether the urokinase cleaving activity is the property of a novel enzyme. activity was assayed by the specific cleavage of recombinant pro-urokinase into 2 fragments, as detd. by SDS-PAGE of unreduced samples of the reaction products of pro-urokinase with the enzyme. The cleaving enzyme was used to prep., in a single digestion, milligram quantities of both an N-terminal fragment of urokinase, comprised of the growth factor and kringle domains, and low-mol.-wt. pro-urokinase. The low-mol.-wt. zymogen could be converted to the active enzyme by careful treatment with plasmin. Low-mol.-wt. urokinase with Leu(144) as its N-terminal residue was produced by culture of human kidney cells. This was in contrast to the low-mol.-wt. urokinase isolated from urine, which begins at Lys (136). Since the N-terminal fragment, Ser(1)-Glu(143), has previously been shown to bind with high affinity to the urokinase receptor, and also

to be active in stimulating the growth or differentiation of certain cells in culture, it is possible this metalloprotease has a role in certain cell

Fibrinolysis (1992), 6(Suppl. 1), 57-62

CODEN: FBRIE7; ISSN: 0268-9499

regulatory functions.

L10 ANSWER 7 OF 8 CAPLUS COPIRIGHT 2002 ACS

TI Development of enzyme-linked immunosorbent assay for free human pro-colipase activation peptide (APGPR)

AΒ Human pancreatic colipase is secreted as the inactive form procolipase. Activation involves tryptic cleavage of an N-terminal pentapeptide Ala-Pro-Gly-Pro-Arg (APGPR) which is known as procolipase activation peptide (CLAP). N-terminally haptenized synthetic APGPR was used to generate specific C-terminally directed anti-APGPR antibodies. The antiserum was used to develop a competitive enzyme linked immunosorbent assay (ELISA) specific for free CLAP with a detection limit of 12 nmol/L and an intra-assay coeff. of variation (CV) of 3.28% and an inter-assay CV of 5.82%. The release of immunoreactive CLAP from human pancreatic juice and chicken pancreas upon trypsinization was demonstrated, as well as the absence of reactivity of the antisera with procolipase from which the CLAP is released. APGPR was found to be unstable in biol. fluids. Immunoreactivity is rapidly lost with half life of 5 min and 4 h in human serum and urine resp. This loss of reactivity can be significantly slowed by the addn. of 20 mmol/L zinc ions (Zn2+), while EDTA and other protease inhibitors were ineffective. In serum the moiety responsible for loss of immunoreactivity was found to have an estd. mol. mass of 200,000-300,000 Da. CLAP assay specifically reports procolipase activation and may help elucidate the mechanism of satiety as well as contribute to the recognition and understanding of the role of procolipase activation in diseases states such as pancreatitis.

SO Clin. Chim. Acta (1991), 200(2-3), 137-52 CODEN: CCATAR; ISSN: 0009-8981

- L10 ANSWER 8 OF 8 CAPLUS COPYRIGHT 2002 ACS
- TI Freeze-dried reagent mixtures for clinical determination of proteases and related factors
- AB A stable, freeze-dried reagent compn. contg., in one container, all substances needed for detn. of a component active in proteolysis is prepd. A soln. contg. acetate buffer, pH 4.2, factor Xa, substrate S-2732, bovine serum albumin, and mannitol was freeze-dried in a plastic cuvet. A plasma sample dild. with a soln. contg. Tris, pH 8.4, EDTA, heparin, and PEG was added to the cuvet. The absorbance at 405 nm was detd. after 8 min incubation and stopping of the reaction with HOAc. The concn. of Factor Xa was detd. by comparing the obsd. absorbance with a std. curve prepd. by the manufacturer of the reagent.

SO PCT Int. Appl., 38 pp.

CODEN: PIXXD2

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L6
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L7
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L8
          83181 S EDTA OR EGTA
L9
           547 S L1 AND L8
L10
              8 S L9 AND URINE
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